

# Expression of a Gibberellin 2-Oxidase Gene around the Shoot Apex Is Related to Phase Transition in Rice<sup>1</sup>

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A major catabolic pathway for gibberellin (GA) is initiated by 2 $\beta$ -hydroxylation, a reaction catalyzed by GA 2-oxidase. We have isolated and characterized a cDNA, designated *Oryza sativa* GA 2-oxidase 1 (*OsGA2ox1*) from rice (*Oryza sativa* L. cv Nipponbare) that encodes a GA 2-oxidase. The encoded protein, produced by heterologous expression in *Escherichia coli*, converted GA<sub>1</sub>, GA<sub>4</sub>, GA<sub>9</sub>, GA<sub>20</sub>, and GA<sub>44</sub> to the corresponding 2 $\beta$ -hydroxylated products GA<sub>8</sub>, GA<sub>34</sub>, GA<sub>51</sub>, GA<sub>29</sub>, and GA<sub>98</sub>, respectively. Ectopic expression of the *OsGA2ox1* cDNA in transgenic rice inhibited stem elongation and the development of reproductive organs. These transgenic plants were deficient in endogenous GA<sub>1</sub>. These results indicate that *OsGA2ox1* encodes a GA 2-oxidase, which is functional not only *in vitro* but also *in vivo*. *OsGA2ox1* was expressed in shoot apex and roots but not in leaves and stems. *In situ* hybridization analysis revealed that *OsGA2ox1* mRNA was localized in a ring at the basal region of leaf primordia and young leaves. This ring-shaped expression around the shoot apex was drastically decreased after the phase transition from vegetative to reproductive growth. It was absent in the floral meristem, but it was still present in the lateral meristem that remained in the vegetative phase. These observations suggest that *OsGA2ox1* controls the level of bioactive GAs in the shoot apical meristem; therefore, reduction in its expression may contribute to the early development of the inflorescence meristem.

Gibberellins (GAs) are endogenous phytohormones that are involved in the regulation of the life cycle of plants. Therefore, biosynthesis of GAs has been intensively studied. Bioactive GAs, such as GA<sub>1</sub> and GA<sub>4</sub>, are synthesized from trans-geranylgeranyl diphosphate by the sequential action of cyclases in plastids, membrane-associated mono-oxygenases in the endoplasmic reticulum, and soluble 2-oxoglutarate-dependent dioxygenases (2ODDs) in the cytosol (Hedden and Kamiya, 1997; Lange, 1998). During the last 10 years, genes for GA 20-oxidase and GA 3 $\beta$ -hydroxylase have been cloned. They encode 2ODDs that catalyze the later steps in GA biosynthesis, namely, the oxidation of the C-20 group and the introduction of the 3 $\beta$ -hydroxyl group, respectively. In *Arabidopsis*, the transcript levels of GA 20-oxidase genes and a GA 3 $\beta$ -hydroxylase gene (GA4) are subject to feedback regulation (Chang et al., 1995; Phillips et al., 1995). These results indicate that the endogenous levels of bioactive GAs are maintained at

proper levels through the regulation of transcript levels of GA 20-oxidase and GA 3 $\beta$ -hydroxylase.

Catabolism of GAs is another important factor that regulates the endogenous levels of bioactive GAs. In many plant species, bioactive GAs are 2 $\beta$ -hydroxylated to produce biologically inactive GAs. This step is catalyzed by a third 2ODD, GA 2-oxidase. This enzyme also inactivates immediate precursors of bioactive GAs such as GA<sub>20</sub> and GA<sub>9</sub> (Ross et al., 1995). Thus, GA 2-oxidase has been considered to play an important role in the regulation of plant growth through the reduction of endogenous levels of bioactive GAs.

The first GA 2-oxidase cDNA was cloned from runner bean (*Phaseolus coccineus*) by functional screening, and then three GA 2-oxidase cDNAs were cloned from *Arabidopsis* by database screening (Thomas et al., 1999). Two cDNAs for GA 2-oxidase were also isolated from garden pea (*Pisum sativum*) by functional screening and reverse transcription (RT)-PCR (Lester et al., 1999; Martin et al., 1999). A loss-of-function mutation in the GA 2-oxidase gene of garden pea results in the slender phenotype. This indicates that GA 2-oxidase has an important role in the regulation of elongation growth (Lester et al., 1999; Martin et al., 1999).

We describe the cloning and characterization of a GA 2-oxidase gene, *OsGA2ox1*, from rice (*Oryza sativa* L. cv Nipponbare). Based on detailed expression analysis of *OsGA2ox1*, we suggest that the GA 2-oxidase

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encoded by this gene participates in the phase transition from vegetative to reproductive growth.

## RESULTS

### Isolation of a GA 2-Oxidase Gene from Rice

The predicted amino acid sequences of conserved regions among 2ODDs, including GA 20-oxidase and GA 3 $\beta$ -hydroxylase (Prescott, 1993), were used to search for putative GA 2-oxidase genes in databases. *Arabidopsis* genomic sequence T31E10.11 (accession no. AC004077) has relatively high homology with the sequences of GA 3 $\beta$ -hydroxylase genes, although its deduced amino acid sequence did not have M-G-L-A-A/P-H-T-D motif that is conserved in the GA 3 $\beta$ -hydroxylases reported. The sequence of *Marah macrocarpa* dioxygenase mRNA M7-3 (accession no. Y09113; MacMillan et al., 1997) had high homology with T31E10.11. The full-length cDNA corresponding to T31E10.11 was isolated from *Arabidopsis* inflorescences by RT-PCR, and the 2 $\beta$ -hydroxylation activity of the recombinant protein was confirmed in vitro. This cDNA was identical to the cDNA clone *AtGA2ox3* reported by Thomas et al. (1999).

The predicted amino acid sequences of *AtGA2ox3*, M7-3, and two rice GA 3 $\beta$ -hydroxylases (H. Itoh, M. Ueguchi-Tanaka, N. Sentoku, H. Kitano, M. Matsuoka, and M. Kobayashi, unpublished data) were compared to design degenerate oligonucleotide primers. Using total RNA from rice shoot as a template, RT-PCR with degenerate primers produced one sequence of the expected length with significant homology to GA 2-oxidases from *Arabidopsis*. This clone was used for further screening to isolate corresponding full-length cDNA and genomic clones.

The isolated full-length cDNA contained an open reading frame of 1,146 bp encoding a protein of 382 amino acids, and was designated *OsGA2ox1* (*Oryza sativa* GA 2-oxidase 1). The predicted amino acid sequence of *OsGA2ox1* contained the amino acids that are conserved within the 2ODDs for GA biosynthesis, including His-241, Asp-243, and His-302 (Fig. 1A, numbers refer to *OsGA2ox1* sequence), the amino acid residues that are supposed to associate at the catalytic site and bind with Fe<sup>2+</sup> (Valegard et al., 1998). When compared with other 2ODDs for GA biosynthesis, the deduced amino acid sequence of *OsGA2ox1* showed highest homology with GA 2-oxidases (Fig. 1B). However, the phylogenetic relationship revealed that GA 2-oxidases from dicot plants share relatively high (47%–69%) amino acid identity with each other but significantly lower (< 44%) identity with *OsGA2ox1* (Fig. 1B; Table I).

The corresponding genomic DNA that completely covers the *OsGA2ox1* coding region was also cloned. By comparing the genomic DNA and cDNA sequences, we revealed that *OsGA2ox1* consists of three exons and two introns (Fig. 1A). This exon/intron structure is also conserved in the *AtGA2ox3* coding sequence.

To investigate the existence of a related sequence for *OsGA2ox1* in the rice genome, we digested rice genomic DNA with several restriction enzymes and subjected it to DNA gel-blot analysis at low stringency, using *OsGA2ox1* cDNA fragment as a probe. As shown in Figure 1C, only one band was obtained; this indicates that the *OsGA2ox1* protein is encoded by a single-copy gene in the rice genome.

### Function of Recombinant *OsGA2ox1* Protein

Recombinant *OsGA2ox1* protein was prepared by expressing the *OsGA2ox1* cDNA in *Escherichia coli*. The catalytic properties of the recombinant protein were examined by incubating cell lysate with GAs identified from rice (Kobayashi et al., 1988, 1989). GAs without a free carboxyl at C-19 were converted to the corresponding 2 $\beta$ -hydroxylated product by the recombinant *OsGA2ox1* protein (Table II). On the other hand, no metabolite was identified for GA<sub>19</sub> and GA<sub>53</sub>; both GAs have a free carboxyl at C-19. This indicates that *OsGA2ox1* is a GA 2-oxidase.

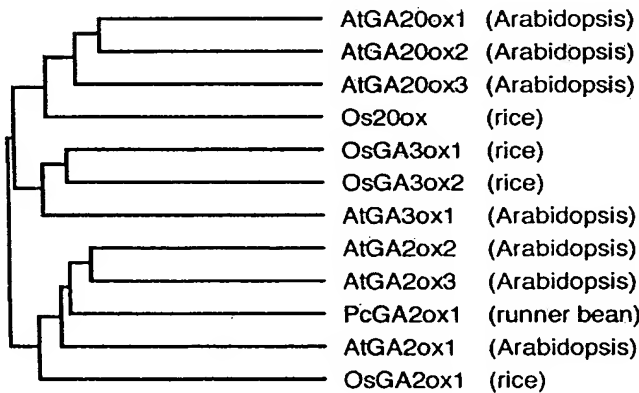
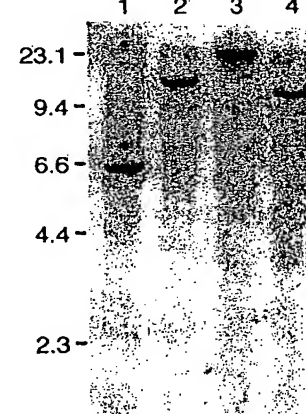
### Ectopic Expression of the *OsGA2ox1* cDNA in Transgenic Rice

To assess the activity of the *OsGA2ox1* gene product in vivo, we fused the full-length *OsGA2ox1* cDNA to the rice actin promoter in the sense orientation and introduced it into wild-type rice by Agrobacterium-mediated gene transfer. All primary transformants (46 independent lines) showed dwarf phenotype. The final plant height of extremely dwarfed transformants was less than 15 cm, whereas that of wild type was 90 cm. Their leaf blades were dark green and shorter and wider than those of wild-type plants, a typical phenotype for GA-deficient dwarf rice (Fig. 2A). Although wild-type plants flowered approximately 90 d after sowing, the formation of floral organs and internode elongation in the extremely dwarfed transformants were not observed even at 120 d after sowing. The transformants did not bear any seeds, but exogenous application of GA<sub>3</sub> could rescue this phenotype (data not shown).

We have previously reported that major endogenous GAs in vegetative tissues of rice are biosynthesized via the early 13-hydroxylation pathway, and GA<sub>1</sub> is the major bioactive GA that regulates shoot elongation (Kobayashi et al., 1988, 1989, 1994). Therefore, we compared the levels of 13-hydroxylated GAs in the leaves of wild-type and extremely dwarfed transgenic plants (Table III). In the transformants, the level of GA<sub>1</sub> decreased to less than one-fourth that of wild-type plants. In contrast, the levels

**A**

OsGA2ox1	MVV-PSATTPARQETVAAAAPPAAASGVGGGGVTTATVDM	SAERGAVARQVATACA	AHGFRCVGHGVPAAPVAA	RLDAATAAFFA	MAPAEK	95													
AtGA2ox1	MAVLSKPVAIPKSGFSL	-----	IPVIDMS	-DPESKRALV	RAKCEDDFGFFKVINHGVS	AE--LVSVLEHETVDP	72												
AtGA2ox2	MVVLPOPVTLDNHSI	LIPTYKPVVLTS	-----	SIPVVNL	A-DEPAKTRIV	RACEEFGPFKVINHGVRP	--LMTRLEQEAI	85											
AtGA2ox3	MVIVLQFASPD	SNLYVNPKCKP	-----	IPVIDLT	-DSDAKTQIV	KACEEFGPFKVINHGVRP	--LLTQLEQEAI	81											
PcGA2ox1	MVVLSPQAL	--NQFP	LLKPFKSTPL	PTG	-----	IPVVDLT	-HFD	AKNLIVNACRDFGFFKL	VINHGVP	LE--LMAN	LENEALRFFK	KSQSEK	81						
PsGA2ox1	MPI	-----	TPSS	-----	S	-----	-KPD	AKTLIV	CEDDFGPKVINHG	VP	LE--AISQ	LESEAFKFFS	LPQTEK	63					
PsGA2ox2	MVV-PSPTSMIRT	KKTKA	-----	-----	-----	VGCIPTIDLSL	ERSQL	SELV	VRA	CEEYGP	VINH	SVP	KPE--VISRL	DEEGIEFFS	76				
OsGA2ox1	QRAGPASPLGYCCRS	ICFGNGDVG	GELEY	YLLL	HANPA	AAVHART	TAIDAM	DPSRFS	SAIV	VNEY	YEA	MKKLACE	I	DLGGLKDPR	YFSKL	TNTNADSD	191		
AtGA2ox1	TQVA-GYPPGYG	N	N	SKRIGRNGDVG	WV	Y	Y	LLM	NN	DSGSGPL	F	PSL	L	KSPGT	PRN	TLSKLVSDQ	NTD	167	
AtGA2ox2	NRAGP	PP	GYG	T	KRIG	PNGDVG	WV	Y	Y	LLM	NN	ANP	Q	LSK	PR	TLS	MLRDE	KSD	181
AtGA2ox3	DKAGP	PP	GYG	T	KRIG	PNGDGL	WV	Y	Y	LLM	NN	ANP	Q	LSK	PR	TLS	MLRDE	KSD	177
PcGA2ox1	DRAGP	PP	GYG	T	KRIG	PNGDGL	WV	Y	Y	LLM	NN	ANP	Q	LSK	PR	TLS	MLRDE	KSD	177
PsGA2ox1	EKAGP	ANP	GYG	T	KRIG	PNGDGL	WV	Y	Y	LLM	NN	ANP	Q	LSK	PR	TLS	MLRDE	KSD	155
PsGA2ox2	RGAGT	PP	GYG	C	KRIG	PNGDGL	WV	Y	Y	LLM	NN	ANP	Q	LSK	PR	TLS	MLRDE	KSD	172
OsGA2ox1	CLLRINH	YHPP	SCNI	H	KLD	HDDQCN	I	KSLV	STK	ASNGGN	I	LMAGGR	I	G	F	YFV	VGD	287	
AtGA2ox1	SILR	LNHY	PP	-----	CPL	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	239	
AtGA2ox2	SCRLR	NHY	PA	-----	A	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	249	
AtGA2ox3	SCRLR	NHY	PA	-----	K	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	244	
PcGA2ox1	SCFRL	NHY	PP	-----	CP	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	246	
PsGA2ox1	CFRLR	NHY	PA	-----	CP	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	224	
PsGA2ox2	SLLR	LNHY	PP	-----	VKKLGNDN	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	251	
OsGA2ox1	VLQALT	TN	GRLI	S	IRH	RV	I	A	T	CR	P	R	L	STI	Y	F	V	G	382
AtGA2ox1	SLQV	MT	NGR	F	SV	R	V	LL	AN	CK	N	MI	Y	FAG	PL	T	Q	V	329
AtGA2ox2	ALQV	MT	NGR	F	SV	K	H	V	LL	AD	T	R	R	I	SMI	Y	FAG	PL	341
AtGA2ox3	TLQV	MT	NGR	F	SV	K	H	V	LL	NT	R	R	I	SMI	Y	FAG	PL	335	
PcGA2ox1	ALQV	MT	NGR	F	SV	K	H	V	LL	AD	T	R	R	I	SMI	Y	FAG	PL	332
PsGA2ox1	SLQV	MT	NGR	F	SV	R	V	LL	AN	G	D	R	R	I	SMI	Y	FAG	PL	309
PsGA2ox2	ALQV	LT	NGR	F	SV	R	V	LL	AN	G	D	R	R	I	SMI	Y	FAG	PL	345

**B****C**

**Figure 1.** Sequence analysis and genomic DNA gel blot analysis of *OsGA2ox1*. **A**, Alignment of deduced amino acid sequences of GA 2-oxidases. Asterisks indicate identical amino acids among the enzymes from various plants. Positions of introns are marked by arrowheads. AtGA2ox1, AtGA2ox2, and AtGA2ox3, GA 2-oxidases from *Arabidopsis* (accession nos. AJ132435, AJ132436, and AJ132437); PcGA2ox1, GA 2-oxidase from runner bean (accession no. A132438); PsGA2ox1 and PsGA2ox2, GA 2-oxidases from garden pea (accession nos. AF100954 and AF100955). **B**, Phylogenetic relationships among GA 2-oxidases. AtGA20ox1, AtGA20ox2, and AtGA20ox3, GA 20-oxidases from *Arabidopsis* (accession nos. X83379, X83380, X83381); Os20ox, GA 20-oxidase from rice (accession no. U50333); AtGA3ox1, GA 3 $\beta$ -hydroxylase from *Arabidopsis* (accession no. L37126); OsGA3ox1 and OsGA3ox2, 3 $\beta$ -hydroxylase from rice (H. Itoh, M. Ueguchi-Tanaka, N. Sentoku, H. Kitano, M. Matsuoka, and M. Kobayashi, unpublished data). **C**, Genomic DNA gel blot analysis of *OsGA2ox1*. Five micrograms of genomic DNA was digested with *Apal* (lane 1), *BamHI* (lane 2), *EcoRI* (lane 3), or *HindIII* (lane 4). *M*, markers are indicated at the left in kilobases.

of GA<sub>8</sub> and GA<sub>29</sub>, the 2 $\beta$ -hydroxylated metabolites of GA<sub>1</sub> and GA<sub>20</sub>, were elevated approximately 2.5- and 6.8-fold, respectively, in the leaves of transformants.

We also determined the effects of exogenous application of GA<sub>1</sub> and GA<sub>3</sub> on the elongation of the second leaf sheath in the transgenic seedlings. GA<sub>1</sub>

can be inactivated by GA 2-oxidase, but GA<sub>3</sub> is not. GA<sub>1</sub> and GA<sub>3</sub> showed an almost equivalent effect on the elongation of dwarf rice, cv Tan-ginbozu (Murakami, 1972). In our results, however, GA<sub>1</sub> was much less active than GA<sub>3</sub> on the elongation of the transformants (Fig. 2B).

**Table I.** Amino acid sequence homology between GA 2-oxidases

Values are percentage of amino acid identity (above the diagonal) or similarity (below the diagonal).							
No.	OsGA2ox1 No. 1	AtGA2ox1 No. 2	AtGA2ox2 No. 3	AtGA2ox3 No. 4	PcGA2ox1 No. 5	PsGA2ox1 No. 6	PsGA2ox2 No. 7
1	—	37	36	36	36	44	38
2	51	—	56	53	57	48	57
3	51	75	—	69	62	47	55
4	49	71	84	—	55	47	53
5	49	74	75	67	—	50	56
6	58	65	65	64	67	—	50
7	52	73	74	68	72	67	—

### Expression of *OsGA2ox1* Gene in Rice

We performed RNA gel-blot analysis to determine the expression pattern of *OsGA2ox1* in various organs of rice. A single strong band was detected in RNA from vegetative shoot apices (Fig. 3A). A relatively weak band was also observed in RNA from inflorescence shoot apices, but not from stems, leaf blades, or rachises.

The expression of *AtGA2ox1* and *AtGA2ox2* is up-regulated by the application of GA<sub>3</sub> (Thomas et al., 1999). We examined the effects of GA<sub>3</sub> and uniconazole (an inhibitor of GA biosynthesis) on the level of *OsGA2ox1* mRNA. We also determined the expression of the rice GA 20-oxidase gene, *Os20ox* (Toomasu et al., 1997), as a control. The expression of *Os20ox* was up-regulated by uniconazole and down-regulated by GA<sub>3</sub> (Fig. 3B), but the levels of *OsGA2ox1* transcripts were not affected (Fig. 3B). Moreover, expression levels of *OsGA2ox1* in the dwarf mutants *d1* (GA-insensitive dwarf; Ueguchi-Tanaka et al., 2000) and *d18* (loss-of-function mutation in the rice GA 3 $\beta$ -hydroxylase gene; Murakami, 1972) were similar to those in the wild-type plants (Fig. 3C).

To determine more precisely the spatial pattern of *OsGA2ox1* expression in rice, we conducted *in situ* hybridization with a digoxigenin-labeled *OsGA2ox1* antisense-strand RNA probe. Figure 4A shows the localization of *OsGA2ox1* mRNA in seedlings. Purple staining, which indicates the presence of *OsGA2ox1* mRNA, occurred around the shoot apical and lateral meristems (arrows) and around the meristems of crown roots in culms (arrowheads). Control sections, hybridized with a sense-strand RNA probe, showed

no signal above background staining (data not shown).

Figure 4B shows a near-median longitudinal section of a rice vegetative shoot apex at a higher magnification than Figure 4A. Lines 1, 2, and 3 indicate approximate planes of the cross-sections shown in Figure 4, E, F, and G, respectively. The *OsGA2ox1* expression appeared as pairs of signals on opposite flanks of the shoot apical meristem (SAM), which was located in the basal region of leaf primordia (Fig. 4B). Serial cross-sections (Fig. 4, E–G) better show the spotted expression of *OsGA2ox1* localized as a doughnut-ring shape around the boundary of leaf primordia (Fig. 4B).

The ring-shaped expression of *OsGA2ox1* around the SAM was drastically decreased just after the phase transition from vegetative to inflorescence stage (Fig. 4C). At the stage of primary rachis-branch primordia differentiation, *OsGA2ox1* was barely expressed around the SAM but was strongly expressed around the lateral meristem, which was still in the vegetative phase (arrow in Fig. 4D).

We also examined the *in situ* mRNA localization of a rice GA 3 $\beta$ -hydroxylase gene, *OsGA3ox2*, around the vegetative shoot apex. As shown in Figure 4H, the *OsGA3ox2* mRNA was strongly expressed in young leaves, but it was not expressed in the SAM or in the basal regions of young leaves throughout the vegetative phase.

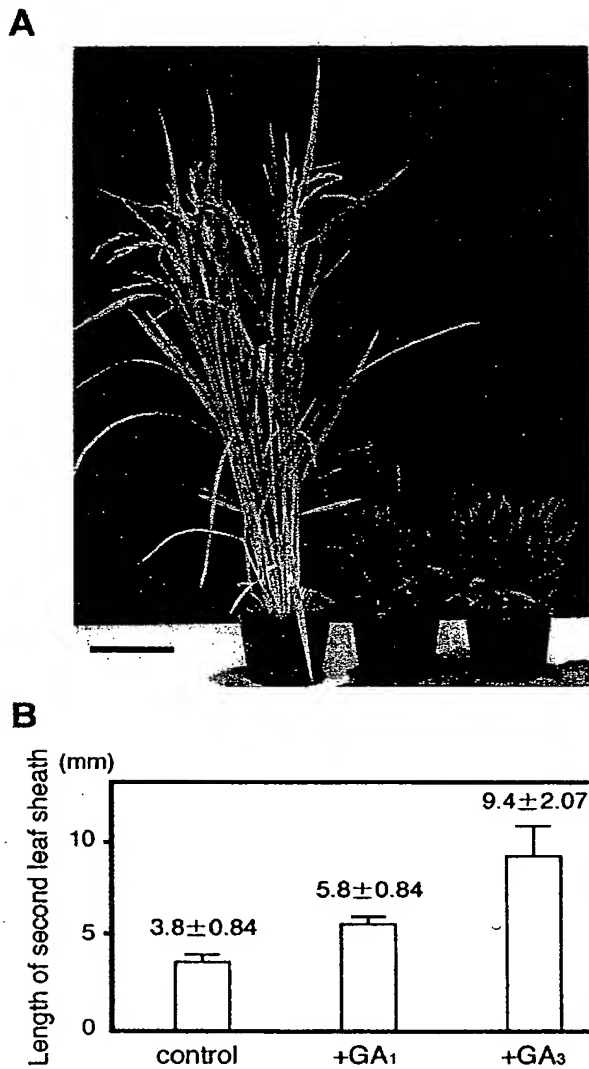
### DISCUSSION

The concentration of bioactive GAs is tightly maintained by the balance between their synthesis and

**Table II.** Metabolism of recombinant *OsGA2ox1* protein

Substrate	Product <sup>a</sup>	KRI	Characteristic Ions at m/z (% Relative Intensity of Base Peak) <sup>b</sup>
<sup>2</sup> H <sub>2</sub> -GA <sub>44</sub>	GA <sub>98</sub>	2,962	522 (47), 507 (5), 373 (4), 345 (4), 240 (21), 209 (100)
<sup>2</sup> H <sub>2</sub> -GA <sub>20</sub>	GA <sub>29</sub>	2,688	509 (100), 494 (6), 450 (2), 379 (6), 210 (25)
<sup>2</sup> H <sub>3</sub> -GA <sub>1</sub>	GA <sub>8</sub>	2,818	597 (100), 582 (7), 538 (8), 451 (5), 210 (20)
<sup>2</sup> H <sub>3</sub> -GA <sub>9</sub>	GA <sub>51</sub>	2,539	389 (22), 331 (30), 287 (78), 271 (76), 227 (100)
GA <sub>4</sub>	GA <sub>34</sub>	2,678	506 (100), 459 (1), 416 (3), 313 (7), 288 (3)

<sup>a</sup> Identification of metabolic products by GC-MS on the basis of KRI and full-scan mass spectra of the methyl ester trimethylsilyl ether derivatives. <sup>b</sup> Based on ions with a mass-to-charge ratio (m/z) of >200.



**Figure 2.** Ectopic expression of *OsGA2ox1* in transgenic rice plants. **A**, Typical phenotype of transgenic rice plants carrying the *Act::OsGA2ox1* gene approximately 120 d after germination. Left, wild-type (cv Nipponbare); center and right, *Act::OsGA2ox1* trans-formants. In contrast to wild-type plants, flowering of transformants was impeded. Bar = 10 cm. **B**, Effect of exogenous GA treatment on elongation of the second leaf sheath of transgenic seedlings. Ten nanograms of GA<sub>1</sub> or GA<sub>3</sub> was dropped on the seedlings, which were then grown for 3 d. Each data point is the average for five plants.

catabolism, as indicated by the fact that GAs reduce the expression of GA20ox genes and stimulate the expression of GA2ox genes. GA 2-oxidase catalyzes the catabolism, namely, the conversion of bioactive GAs and their immediate precursors into biologically inactive GAs by 2 $\beta$ -hydroxylation (Ross et al., 1995). The expression of GA 2-oxidase genes from *Arabidopsis* was stimulated by the application of GA<sub>3</sub> (Thomas et al., 1999). This indicates that the expression of these genes is regulated through feedback to maintain endogenous levels of bioactive GAs.

We cloned *OsGA2ox1*, the first GA 2-oxidase gene from rice. Four of six GA 2-oxidases from runner bean, *Arabidopsis*, and garden pea catalyze not only 2 $\beta$ -hydroxylation but also the further oxidation at C-2. On the other hand, the recombinant *OsGA2ox1* protein catalyzed a single step of the 2 $\beta$ -hydroxylation. These results demonstrate that *OsGA2ox1* inactivates the bioactive GAs and their precursors. The physiological and biochemical phenotypes of transgenic rice plants that ectopically express the *OsGA2ox1* cDNA further support this conclusion.

Two or more GA 2-oxidase genes have been cloned from *Arabidopsis* and garden pea, and their expression was observed in various organs (Lester et al., 1999; Martin et al., 1999; Thomas et al., 1999). Because *OsGA2ox1* was expressed only in limited tissues and was not induced by the application of GA<sub>3</sub>, there may be another GA 2-oxidase gene that maintains bioactive GA levels in various organs of rice. We have tried unsuccessfully so far to clone another GA 2-oxidase gene from rice. Only one band appeared in the DNA gel blot analysis (Fig. 2C). Another GA 2-oxidase gene may have relatively less sequence similarity with those genes already reported. Our recent results have revealed that one of the rice 3 $\beta$ -hydroxylases, *OsGA3ox1*, had activity for 2 $\beta$ -hydroxylation of bioactive GAs (H. Itoh, M. Ueguchi-Tanaka, N. Sentoku, H. Kitano, M. Matsuoka, and M. Kobayashi, unpublished data). Thus, there are at least two genes in the rice genome whose products have 2 $\beta$ -hydroxylation activity.

*OsGA2ox1* was expressed in a limited region around the shoot apex and was not affected by the levels of bioactive GAs. Tissue-specific expression has been reported for *GA4H*, a GA 3 $\beta$ -hydroxylase gene expressed in germinating seeds of *Arabidopsis*. The expression of *GA4H* was induced by the phytochrome signal to promote germination and was not affected by the exogenous application of GA<sub>3</sub> or uniconazole (Yamaguchi et al., 1999). *OsGA2ox1*, similarly, may have a specific role in the regulation of developmental processes in the shoot apex, rather than in the maintenance of bioactive GA levels in leaf tissues.

It is noteworthy that *OsGA3ox2*, the major GA 3 $\beta$ -hydroxylase in the vegetative tissue of rice (H. Itoh, M. Ueguchi-Tanaka, N. Sentoku, H. Kitano, M. Matsuoka, and M. Kobayashi, unpublished data), was expressed in young leaves but not in the SAM itself (Fig. 4H). This result is consistent with the previous report that the young leaves are the source of bioactive GAs (Choi et al., 1995). In contrast, the *OsGA2ox1* mRNA was localized in a ring at the basal region of young leaves. This pattern of gene expression leads us to speculate that *OsGA2ox1* inactivates GAs synthesized in young leaves to keep the levels of bioactive GAs low in the SAM. This may be one of

**Table III.** Concentration (ng g<sup>-1</sup> fresh weight) of GAs in leaves of wild-type and transgenic rice plant overexpressing *OsGA2ox1* cDNA

The level of endogenous GAs in the shoots of wild-type (cv Nipponbare) and transgenic plants were analyzed by GC-SIM using internal standards.

Plant	GA <sub>53</sub>	GA <sub>44</sub>	GA <sub>19</sub>	GA <sub>20</sub>	GA <sub>1</sub>	GA <sub>8</sub>	GA <sub>29</sub>
Wild type	0.37	0.41	2.72	0.66	0.21	0.39	0.17
Transformant	0.20	0.31	5.97	0.74	0.05	0.97	1.15

the mechanisms for repressing internode elongation during the vegetative phase.

It is interesting that the expression of *OsGA2ox1* around the shoot apex disappeared after flower induction. It has been suggested that GA is involved in the phase transition in various plant species (Evans and Poethig, 1995; Chien and Sussex, 1996; McDaniel and Hartnett, 1996; Telfer et al., 1997; Scott et al., 1999). In a long-day plant, *Arabidopsis*, GA has been proposed to promote flowering by activating the *LEAFY* promoter (Okamuro et al., 1996; Blazquez et al., 1997, 1998; Blazquez and Weigel, 1999, 2000). GA also activates *SA-MADS A*, a gene involved in the regulation of floral transition in the long-day plant *Sinapis alba* (Bonhomme et al., 2000). In a short-day

plant, *Pharbitis nil*, application of uniconazole inhibited the flowering response induced by short-day treatment, and the inhibition by uniconazole was canceled by further application of GA<sub>1</sub> (Wijayanti et al., 1996, 1997). These studies indicate that GA is required for the differentiation and/or development of the shoot apex during the phase transition. This hypothesis is consistent with the fact that, in transgenic rice plants ectopically expressing the *OsGA2ox1* cDNA, flowering time was significantly delayed and the development of reproductive organs was impeded. From this point of view, the down-regulation of *OsGA2ox1* in the shoot apex may be one of the initial events required for the development of inflorescence meristem.

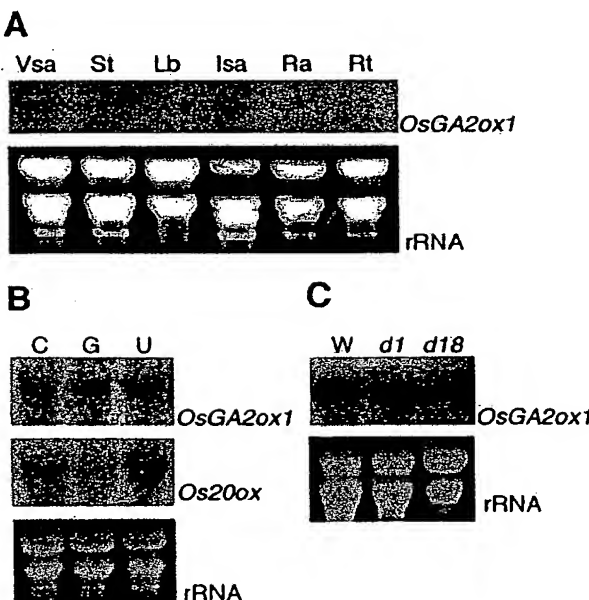
## MATERIALS AND METHODS

### Plant Materials

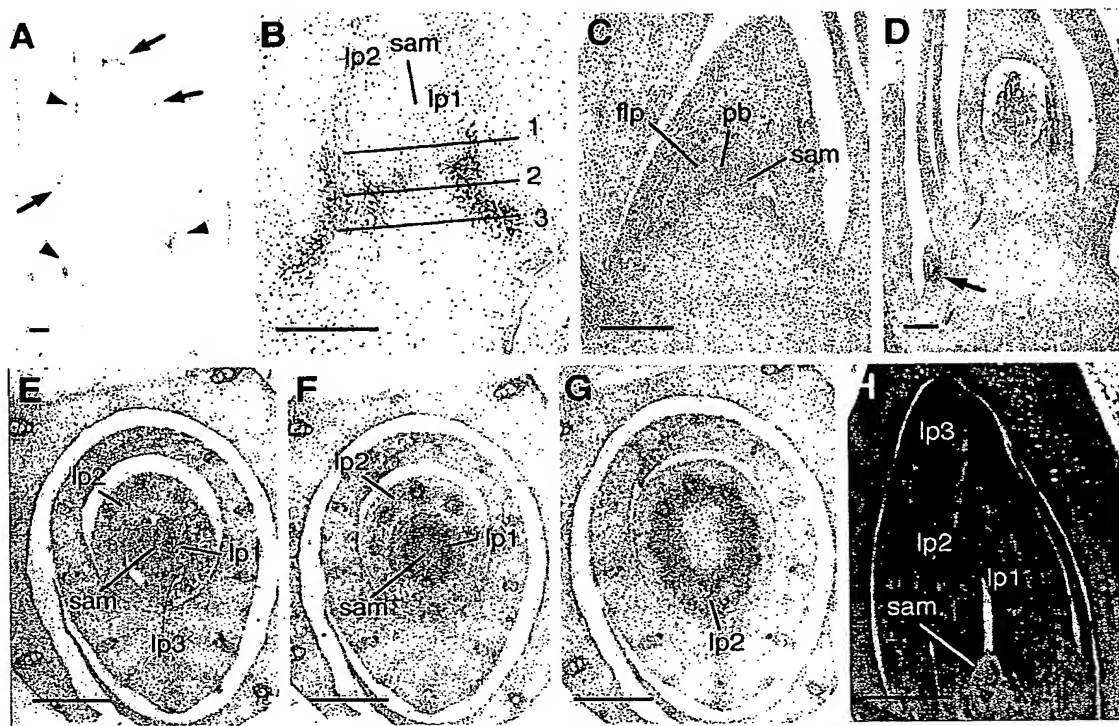
Seeds of tall rice (*Oryza sativa* L. cv Nipponbare), two GA-insensitive dwarf cultivars (Daikoku-dwarf, *T65-d1*, and Hosetsu-dwarf, *d18-ID18h*), and *OsGA2ox1* transformants were sterilized in 1% (v/v) NaClO for 1 h and sown on an agar medium. Seedlings were grown in a growth chamber with a cycle of 16-h light/8-h dark at 30°C. To investigate the effects of GA<sub>3</sub> and uniconazole on the expression of *OsGA2ox1*, seeds of wild-type rice were sown on an agar medium containing 10 μM GA<sub>3</sub> or 10 μM uniconazole and grown for 3 d.

### Isolation of Partial Fragment Encoding Rice GA 2-Oxidase

To amplify a GA 2-oxidase from rice, total RNA was extracted from whole seedlings. RT-PCR was performed by using two degenerate oligonucleotide primers (forward primer, 5'-GGNTTYGGNGARCAWCNGAYCC-3'; reverse primer, 5'-ACNCKRTGYYTAYISWIWYVARICKICCRTT-GT-3'). Amplified fragments (approximately 200 bp) were cloned into pCR II (Invitrogen, Carlsbad, CA), and their sequences were determined. One of the 64 independent clones showed homology with 2ODDs and was predicted to encode a rice GA 2-oxidase gene. The deduced amino acid sequence for this clone was used to search the DDBJ nucleotide sequence database, and one rice expressed sequence tag clone (accession no. C72618) was obtained. Oligonucleotide primers were designed based on the sequence of this clone (forward primer, 5'-GCGCGTTCTTCGCG-3'; reverse primer, 5'-ATGCTAGCCTTGGTGTCAA-3') and used in RT-PCR. An amplified fragment (230 bp) was cloned into



**Figure 3.** RNA gel-blot analysis of *OsGA2ox1*. Ten micrograms of total RNA was loaded per lane. The blot was probed with <sup>32</sup>P-labeled *OsGA2ox1* or *Os20ox* cDNA. The lower part of A and B shows ethidium bromide-stained RNAs corresponding to the above lanes for loading control. A, *OsGA2ox1* expression in various organs of wild-type rice. Total RNA was extracted from vegetative shoot apices (Vsa), stems (St), leaf blades (Lb), inflorescence shoot apices (Isa), rachises (Ra), and roots (Rt). B, Effect of GA<sub>3</sub> and uniconazole treatment on the levels of *OsGA2ox1* and *Os20ox* transcripts. Total RNA was extracted from wild-type seedlings treated with 10 μM GA<sub>3</sub> (G) or 10 μM uniconazole (U) or from untreated control plants (C). C, *OsGA2ox1* expression in rice dwarf mutants. Total RNA was extracted from wild-type (W), GA-insensitive dwarf mutant (*d1*), and GA-responsive dwarf mutant (*d18*) seedlings.



**Figure 4.** *In situ* mRNA localization of *OsGA2ox1* and *OsGA3ox2*. Purple staining indicates the presence of *OsGA2ox1* (A–G) and *OsGA3ox2* (H) mRNA. A, Longitudinal section of seedling at 7 d after germination. Arrows and arrowheads indicate the shoot and root meristem, respectively. B, Median longitudinal section of the vegetative shoot apex at higher magnification than (A). Lines 1, 2, and 3 indicate the approximate planes of bisection shown in E, F, and G. C, Median longitudinal section of the inflorescence meristem. D, Median longitudinal section of the floral meristem at the primary rachis branch stage. Arrow indicates the lateral meristem. E through G, Serial cross sections around the vegetative shoot apex. H, Longitudinal section of seedling at 3 d after sowing. sam, Shoot apical meristem; lp1 through 3, leaf primordium 1 through 3; flp, flag leaf primordium; pb, primary rachis branch.

pCR II (Invitrogen), and the sequence was confirmed. This DNA fragment was used to screen the cDNA and genomic libraries.

#### Screening of Rice cDNA and Genomic Libraries

A cDNA library was constructed with mRNA prepared from immature rice seed, and a genomic library was constructed from rice genomic DNA partly digested with *Sau3AI*. The libraries were screened with a  $^{32}$ P-labeled probe prepared as described above. Hybridization was performed in 5× SSC (1× SSC contains 0.15 M NaCl, 15 mM sodium citrate), 5× Denhardt's solution (1× Denhardt's solution contains 0.02% [w/v] Ficoll, 0.02% [w/v] polyvinylpyrrolidone, 0.02% [w/v] bovine serum albumin), 0.5% (w/v) SDS, and 20 mg L<sup>-1</sup> salmon sperm DNA at 65°C for 14 h. Then filters were washed in 2× SSC, 0.1% (w/v) SDS, at room temperature.

#### Sequence Analysis

Nucleotide sequences were determined by a dideoxynucleotide chain-termination method using an automated sequencing system (ABI377). The cDNA and genomic clones were completely sequenced on both strands, includ-

ing a large intron. The cDNA and amino acid sequences were analyzed by Lasergene computer software (DNAStar, Madison, WI).

#### DNA Gel-Blot Analysis

Five micrograms of the genomic DNA isolated from rice leaves was digested with *Apal*, *BamHI*, *EcoRI*, and *HindIII*, electrophoresed on a 0.7% (w/v) agarose gel, then transferred on to nylon membranes (Hybond N<sup>+</sup> membrane Amersham, Buckinghamshire, England) and hybridized with a  $^{32}$ P-labeled probe prepared as described above. Hybridization was performed in 5× SSC, 5× Denhardt's solution, 0.5% (w/v) SDS, 10% (w/v) dextran sulfate, and 20 mg L<sup>-1</sup> salmon sperm DNA at 65°C for 14 h. The filter was washed in 2× SSC, 0.1% (w/v) SDS, at 65°C, and then further washed in 0.2× SSC, 0.1% (w/v) SDS, at 65°C.

#### Enzyme Assays

Full-length *OsGA2ox1* cDNA was inserted in the sense orientation as a translational fusion into the pMAL-c2 expression vector (New England Biolabs, Beverly, MA), and expressed in *Escherichia coli* strain JM109. Cell lysate was prepared as described by Rebers et al. (1999). The presence

of OsGA2ox1 recombinant protein in the cell lysate was confirmed by SDS-PAGE. After the addition of 2-ketoglutarate (final concentration, 5 mM), ascorbate (5 mM), and  $\text{FeSO}_4$  (0.5 mM), the cell lysate (total volume, 200  $\mu\text{L}$ ) was incubated with substrate GA at 30°C for 1 h. Then the reaction was adjusted to pH 2, and the product was extracted with ethyl acetate. The extract was analyzed by full-scan gas chromatography-mass spectrometry with an Automass mass spectrometer (JEOL, Akishima, Japan) connected to a Hewlett-Packard 5890 series II gas chromatograph (Kobayashi et al., 1996). Product GAs were identified by their mass spectrum and Kovats retention index.

#### Plasmid Constructs and Plant Transformation

The full-length cDNA of *OsGA2ox1* was excised as an *Xba*I-*Eco*RV fragment and inserted between the rice actin promoter and the nopaline synthase polyadenylation signal of hygromycin-resistant binary vector pAct-Hm2. This vector is modified from pBI-H1 (Ohta et al., 1990) and contains a rice actin promoter. The resulting fusion construct was introduced into *Agrobacterium tumefaciens* strain EHA101 by electroporation. *Agrobacterium*-mediated transformation of rice was performed as described by Hiei et al. (1994). Transgenic plants were selected on media containing 50 mg  $\text{L}^{-1}$  hygromycin.

#### Analysis of Endogenous GA levels

Endogenous GA levels in the transgenic plants were analyzed by a modification of the procedure of Choi et al. (1995). Plant tissues were homogenized in liquid nitrogen and extracted twice with an excess volume of 80% (v/v) methanol. The extract was amended with 5 ng of deuterated GAs ( $^2\text{H}_2\text{-GA}_{53, 44, 19, 20, 29, 8}$  and  $^2\text{H}_5\text{-GA}_1$ ) as internal standards. After concentration in vacuo, the aqueous residue was submitted to a solvent fractionation procedure to give the acidic ethyl acetate-soluble fraction. After two steps of purification with Bond Elut C<sub>18</sub> and Bond Elut DEA cartridge columns (Varian, Harbor City, CA; Kobayashi et al., 1993), the fraction was submitted to HPLC on a column of Nucleosil 5 C<sub>18</sub> (i.d., 6 mm; length, 100 mm; Macherey-Nagel, Düren, Germany). The column was eluted with a mixture of solvents, A (10% [v/v] methanol, 0.05% [v/v] acetic acid) and B (0.05% [v/v] acetic acid in methanol). After injection, the concentration of solvent B was increased from 20% to 75% over 32 min. The elute was collected according to the retention time of each GA, concentrated, and derivatized to methyl ester trimethylsilyl ether or trimethylsilyl ester trimethylsilyl ether. It was then analyzed by gas chromatography-selected ion monitoring (GC-SIM). The conditions for GC-SIM were as described by Kobayashi et al. (1996).

#### Biological Activity of GA<sub>1</sub> and GA<sub>3</sub> on the Elongation of *OsGA2ox1* Transgenic Plants

The biological activity was evaluated by micro-drop assay (Murakami, 1972).

#### RNA Gel-Blot Analysis

Total RNA was separately prepared from various organs of rice for analysis. Ten micrograms of each RNA preparation was electrophoresed on a 1.2% (w/v) agarose gel, then transferred on to Hybond N<sup>+</sup> membrane and hybridized with a <sup>32</sup>P-labeled probe prepared as described above. The hybridization and washing conditions were as for the DNA gel-blot analysis.

#### In Situ Hybridization Analysis

Plant materials were fixed in 4% (w/v) paraformaldehyde and 0.25% (w/v) glutaraldehyde in 100 mM sodium phosphate buffer (pH 7.4) overnight at 4°C, dehydrated through a graded ethanol series and *t*-butanol series (Sass, 1958), and finally embedded in Paraplast Plus (Sherwood Medical, St. Louis). Microtome sections (7–10  $\mu\text{m}$  thick) were applied to glass slides treated with silane. Hybridization with a digoxigenin-labeled RNA probe and immunological detection were conducted according to the methods of Kouchi and Hata (1993).

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